```
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TERMINAL (ENTER 1, 2, 3, OR ?):2

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                Zentralblatt
NEWS 3 OCT 19 BEILSTEIN updated with new compounds
NEWS 4 NOV 15 Derwent Indian patent publication number format enhanced
NEWS 5 NOV 19 WPIX enhanced with XML display format
NEWS 6 NOV 30 ICSD reloaded with enhancements
NEWS 7 DEC 04 LINPADOCDB now available on STN
NEWS 8 DEC 14 BEILSTEIN pricing structure to change
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NEWS 10 DEC 17 IMSDRUGCONF removed from database clusters and STN
NEWS 11 DEC 17 DGENE now includes more than 10 million sequences
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                MEDLINE segment
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NEWS 17 JAN 16 CAS patent coverage enhanced to include exemplified
                prophetic substances
NEWS 18 JAN 28 USPATFULL, USPAT2, and USPATOLD enhanced with new
                custom IPC display formats
NEWS 19 JAN 28 MARPAT searching enhanced
NEWS 20 JAN 28 USGENE now provides USPTO sequence data within 3 days
                of publication
NEWS 21 JAN 28 TOXCENTER enhanced with reloaded MEDLINE segment
NEWS 22 JAN 28 MEDLINE and LMEDLINE reloaded with enhancements
NEWS 23 FEB 08 STN Express, Version 8.3, now available
NEWS 24 FEB 20 PCI now available as a replacement to DPCI
NEWS 25 FEB 25 IFIREF reloaded with enhancements
NEWS 26 FEB 25 IMSPRODUCT reloaded with enhancements
NEWS 27 FEB 29 WPINDEX/WPIDS/WPIX enhanced with ECLA and current
                U.S. National Patent Classification
NEWS EXPRESS FEBRUARY 08 CURRENT WINDOWS VERSION IS V8.3.
            AND CURRENT DISCOVER FILE IS DATED 20 FEBRUARY 2008
```

NEWS HOURS STN Operating Hours Plus Help Desk Availability

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FILE 'HOME' ENTERED AT 18:40:49 ON 11 MAR 2008

=> fil req

FULL ESTIMATED COST

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION

0.21

0.21

FILE 'REGISTRY' ENTERED AT 18:41:03 ON 11 MAR 2008

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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 10 MAR 2008 HIGHEST RN 1007341-18-5 DICTIONARY FILE UPDATES: 10 MAR 2008 HIGHEST RN 1007341-18-5

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH January 9, 2008.

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

Uploading C:\Program Files\Stnexp\Queries\10507925.str

chain nodes : 10 11 12 ring nodes :

1 2 3 4 5 6 7 8 9 13 14 15 16 17 18 19 20 21 chain bonds :

1-10 10-11 11-12 12-13

ring bonds :

1-2 1-6 2-3 3-4 3-7 4-5 4-9 5-6 7-8 8-9 13-14 13-18 14-15 15-16 16-17

16-19 17-18 17-21 19-20 20-21 exact/norm bonds: 1-10 3-7 4-9 7-8 8-9 10-11 11-12 12-13 16-19 17-21 19-20 20-21 normalized bonds: 1-2 1-6 2-3 3-4 4-5 5-6 13-14 13-18 14-15 15-16 16-17 17-18

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:CLASS 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:Atom 20:Atom 21:Atom

L1 STRUCTURE UPLOADED

=> d 11 L1 HAS NO ANSWERS L1 STR

Structure attributes must be viewed using STN Express query preparation.

=> s ss 11 sam SAMPLE SEARCH INITIATED 18:41:29 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 1280 TO ITERATE

100.0% PROCESSED 1280 ITERATIONS 1 ANSWERS SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 23454 TO 2705
PROJECTED ANSWERS: 1 TO 80

L2 1 SEA SSS SAM L1

=> d scan

L2 1 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN 2-Benzofurancarboxylic acid, 6,6'-[methylenebis(oxy)]bis[3-methyl-, diethyl ester (9CI)

MF C25 H24 O8

$$\begin{array}{c} \text{Me} \\ \text{Et} \ O \\ \end{array} \begin{array}{c} O \\ \text{CH} \ 2 \\ \end{array} \begin{array}{c} O \\ \text{CH} \ 2 \\ \end{array} \begin{array}{c} O \\ \text{Et} \\ \end{array} \begin{array}{c} O \\ \text{Et} \\ \end{array} \begin{array}{c} O \\ \text{Et} \\ \end{array}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ALL ANSWERS HAVE BEEN SCANNED

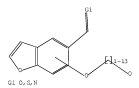
=>

Uploading C:\Program Files\Stnexp\Queries\10507925B.str

```
chain nodes:
10 11 12 15 16
ring nodes:
1 2 3 4 5 6 7 8 9
chain bonds:
6-15 10-11 11-12 15-16
ring bonds:
1-2 1-6 2-3 3-4 3-7 4-5 4-9 5-6 7-8 8-9
exact/norm bonds:
3-7 4-9 7-8 8-9 10-11 11-12 15-16
exact bonds:
6-15 normalized bonds:
1-2 1-6 2-3 3-4 4-5 5-6
```

G1:0,S,N

Match level: 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 15:CLASS 15:CLASS 16:CLASS 18:Atom



Structure attributes must be viewed using STN Express query preparation.

1 ANSWERS

=> S SSS L3 SAM

SAMPLE SEARCH INITIATED 18:55:45 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 14700 TO ITERATE

13.6% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 286737 TO 301263

PROJECTED ANSWERS: 1 TO

L4 1 SEA SSS SAM L3

=> D SCAN

L4 1 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN Ethanone, 1-[6-hydroxy-4-methoxy-7-(3-phenoxypropoxy)-5-benzofurany1]-

309

MF C20 H20 O6

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ALL ANSWERS HAVE BEEN SCANNED

=>

Uploading C:\Program Files\Stnexp\Queries\10507925C.str

chain nodes:
10 11 12 15 16
ring nodes:
1 2 3 4 5 6 7 8 9
chain bonds:
6-15 10-11 11-12 15-16
ring bonds:
1-2 1-6 2-3 3-4 3-7 4-5 4-9 5-6 7-8 8-9
exact/norm bonds:
3-7 4-9 7-8 8-9 10-11 11-12 15-16
exact bonds:
6-15 normalized bonds:
1-2 1-6 2-3 3-4 4-5 5-6

G1:0,S,N

Match level: 1:1Atom 2:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:CLASS 15:CLASS 16:CLASS 18:Atom

L5 STRUCTURE UPLOADED

=> D L5 L5 HAS NO ANSWERS L5 STR

Structure attributes must be viewed using STN Express query preparation.

1 ANSWERS

=> S SSS L5 SAM

SAMPLE SEARCH INITIATED 18:58:39 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 14700 TO ITERATE

13.6% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE**

PROJECTED ITERATIONS: 286737 TO 301263 PROJECTED ANSWERS: 1 TO 309

L6 1 SEA SSS SAM L5

=> D SCAN

L6 1 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN Ethanone, 1-[6-hydroxy-4-methoxy-7-(3-phenoxypropoxy)-5-benzofurany1]-MF

C20 H20 O6

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ALL ANSWERS HAVE BEEN SCANNED

=> S SSS L5 FULL

FULL SEARCH INITIATED 18:58:53 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 292385 TO ITERATE

100.0% PROCESSED 292385 ITERATIONS SEARCH TIME: 00.00.02 125 ANSWERS

L7 125 SEA SSS FUL L5

=> D SCAN

L7 125 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN Acetic acid, [[4,7-dimethoxy-5-[1-oxo-3-(3,4,5-trimethoxyphenyl)-2-propenyl]-6-benzofuranyl]oxy]- (9CI)

MF C24 H24 O10

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):4

- L7 125 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN Acetic acid, [(5-acetyl-6-hydroxy-7-methoxy-4-benzofuranyl)oxy]- (6CI, 9CI)
- MF C13 H12 O7

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L7 125 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN Ethanone, 1,1'-[1,5-pentanediylbis[oxy(4,7-dimethoxy-6,5-benzofurandiyl)]]bis- (9CI)
- MF C29 H32 O10

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L7 125 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN Ethanone, 1-[7-[2-(4-fluorophenoxy)ethoxy]-6-hydroxy-4-methoxy-5benzofurany1]-
- MF C19 H17 F 06

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L7 125 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN Carbonic acid, ethyl 5-[[(4-hydroxyphenyl)amino]carbonyl]-4-methoxy-6benzofuranyl ester (9CI)
- MF C19 H17 N 07

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> SAVE TEMP L7 BAEL/10507925/A

BAEL/10507925/A IS NOT A VALID SAVED NAME

Enter the name you wish to use for the saved query, answer set, or L-number list. The name must:

- 1. Begin with a letter,
- 2. Have 1-12 characters,
- 3. Contain only letters (A-Z) and numbers (0-9),
- End with /Q for a query (search profile, structure, or screen set), /A for an answer set, or /L for an L-number list.
- 5. Not already be in use as a saved name,
- 6. Not be END, SAV, SAVE, SAVED
- 7. Not have the form of an L-number (Lnnn). ENTER NAME OR (END):END

ANSWER SET L7 HAS BEEN SAVED AS 'BAEL10507925/A'

=> SAVE TEMP L7 BAEL10507925/A

=>

Uploading C:\Program Files\Stnexp\Queries\10507925D.str

chain nodes :

10 11 12 15 16 27 28

ring nodes:
1 2 3 4 5 6 7 8 9 18 19 20 21 22 23 24 25 26

chain bonds : 1-10 6-15 10-11 11-12 12-18 15-16 19-27 27-28

ring bonds :

 $exact/norm\ bonds$:

1-10 3-7 4-9 7-8 8-9 10-11 11-12 12-18 15-16 21-24 22-26 24-25 25-26 27-28

exact bonds :

6-15 19-27

normalized bonds :

 $1-2 \quad 1-6 \quad 2-3 \quad 3-4 \quad 4-5 \quad 5-6 \quad 18-19 \quad 18-23 \quad 19-20 \quad 20-21 \quad 21-22 \quad 22-23$

G1:0, S, N

Match level:
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS
11:CLASS 12:CLASS 15:CLASS 16:CLASS 18:Atom 19:Atom 20:Atom 21:Atom 22:Atom 23:Atom 24:Atom 25:Atom 26:Atom 27:CLASS 28:CLASS

L8 STRUCTURE UPLOADED

=> D L8 L8 HAS NO ANSWERS L8 STR

Structure attributes must be viewed using STN Express query preparation.

=> S SSS L8 SAM

SAMPLE SEARCH INITIATED 19:05:47 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 94 TO ITERATE

100.0% PROCESSED 94 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE** PROJECTED ITERATIONS: 1299 TO 246

PROJECTED ITERATIONS: 1299 TO 2461 PROJECTED ANSWERS: 0 TO 0

L9 0 SEA SSS SAM L8

=> S SSS L8 SUBSET=L7 SAM

SAMPLE SUBSET SEARCH INITIATED 19:06:23 FILE 'REGISTRY'
SAMPLE SUBSET SCREEN SEARCH COMPLETED - 0 TO ITERATE

100.0% PROCESSED 0 ITERATIONS 0 ANSWERS SEARCH TIME: 00.00.01

PROJECTIONS (WITHIN SPECIFIED SUBSET): ONLINE **COMPLETE**
PROJECTED ITERATIONS (WITHIN SPECIFIED SUBSET): 0 TO 0
PROJECTED ANSWERS (WITHIN SPECIFIED SUBSET): 0 TO 0

L10 0 SEA SUB=L7 SSS SAM L8

=> D SCAN L10 HAS NO ANSWERS

=>

Uploading C:\Program Files\Stnexp\Queries\10507925E.str

chain nodes:
10 11 12 15 16 27 28

ring nodes:
1 2 3 4 5 6 7 8 9 18 19 20 21 22 23 24 25 26

chain bonds:
1-10 6-15 10-11 11-12 12-18 15-16 19-27 27-28

ring bonds:
1-2 1-6 2-3 3-4 3-7 4-5 4-9 5-6 7-8 8-9 18-19 18-23 19-20 20-21 21-22

21-24 22-23 22-26 24-25 25-26

exact/norm bonds:
1-10 3-7 4-9 7-8 8-9 10-11 11-12 12-18 15-16 21-24 22-26 24-25 25-26

exact bonds:
27-28

exact bonds:
6-15 19-27

1-2 1-6 2-3 3-4 4-5 5-6 18-19 18-23 19-20 20-21 21-22 22-23

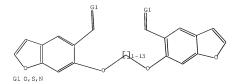
G1:0,S,N

normalized bonds :

Match level: 1: 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:CLASS 15:CLASS 16:CLASS 18:Atom 19:Atom 20:Atom 21:Atom 22:Atom 23:Atom 24:Atom 25:Atom 26:Atom 27:CLASS 28:CLASS

L11 STRUCTURE UPLOADED

=> D L11 L11 HAS NO ANSWERS L11 STR



Structure attributes must be viewed using STN Express query preparation.

=> S SSS L11 SAM SAMPLE SEARCH INITIATED 19:08:16 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 112 TO ITERATE

100.0% PROCESSED 112 ITERATIONS SEARCH TIME: 00.00.01

0 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE** PROJECTED ITERATIONS: 1606 TO 2874 PROJECTED ANSWERS: 0 TO

L12 0 SEA SSS SAM L11

=> D HIS

(FILE 'HOME' ENTERED AT 18:40:49 ON 11 MAR 2008)

FILE 'REGISTRY' ENTERED AT 18:41:03 ON 11 MAR 2008 L1 STRUCTURE UPLOADED L2 1 S SSS L1 SAM STRUCTURE UPLOADED L3 L4 1 S SSS L3 SAM L5 STRUCTURE UPLOADED L6 1 S SSS L5 SAM L7 125 S SSS L5 FULL SAVE TEMP L7 BAEL10507925/A 1.8 STRUCTURE UPLOADED T. 9 0 S SSS L8 SAM L10 0 S SSS L8 SAM SUB=L7 L11 STRUCTURE UPLOADED L12 0 S SSS L11 SAM

=> S SSS L11 SUBSET=L7 SAM

SAMPLE SUBSET SEARCH INITIATED 19:09:07 FILE 'REGISTRY' SAMPLE SUBSET SCREEN SEARCH COMPLETED - 0 TO ITERATE

100.0% PROCESSED 0 ITERATIONS 0 ANSWERS SEARCH TIME: 00.00.01

PROJECTIONS (WITHIN SPECIFIED SUBSET): ONLINE **COMPLETE**

0 TO PROJECTED ITERATIONS (WITHIN SPECIFIED SUBSET): PROJECTED ANSWERS (WITHIN SPECIFIED SUBSET): 0 TO

0 SEA SUB=L7 SSS SAM L11

=> S SSS L11 SUBSET=L7 FULL FULL SUBSET SEARCH INITIATED 19:09:24 FILE 'REGISTRY' FULL SUBSET SCREEN SEARCH COMPLETED - 17 TO ITERATE

100.0% PROCESSED 17 ITERATIONS 5 ANSWERS

SEARCH TIME: 00.00.01

T.14 5 SEA SUB=L7 SSS FUL L11

=> FIL CAPL

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 241.62 241.83

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FILE COVERS 1907 - 11 Mar 2008 VOL 148 ISS 11 FILE LAST UPDATED: 10 Mar 2008 (20080310/ED)

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=> S L14 L15

4 L14

=> D IBIB ABS HITSTR 1-4

L15 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2006:87666 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 144:331145

A New Class of Blockers of the Voltage-Gated Potassium TITLE: Channel Kv1.3 via Modification of the 4- or 7-Position

of Khellinone

AUTHOR(S): Harvey, Andrew J.; Baell, Jonathan B.; Toovey, Nathan;

Homerick, Daniel; Wulff, Heike

CORPORATE SOURCE: The Walter and Eliza Hall Institute, Medical Research Biotechnology Centre, Bundoora, 3086, Australia

SOURCE: Journal of Medicinal Chemistry (2006), 49(4),

1433-1441

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER: American Chemical Society
DOCUMENT TYPE: Journal

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 144:331145

AB The voltage-gated potassium channel Kvl.3 constitutes an attractive target for the selective suppression of effector memory T cells in autoimmune diseases.

We have previously reported the natural product khellinone, as a versatile lead mol. and identified two new classes of Kv1.3 blockers: (1) chalcone derivs. of khellinone, and (ii) khellinone dimers linked through the 6-position. Here we describe the multiple parallel synthesis of a new class of khellinone derivs. selectively alkylated at either the 4- or 7-position via the phenolic OH and show that several chloro, bromo, methoxy, and nitro substituted benzyl derivs. inhibit Kv1.3 with submicromolar potencies. Representative examples of the most potent compds. from each subclass, (5-

AspleseMattive examples of the most potent compus. From each subclass, (3-deety1-4-(4'-chloro)benzyloxy-6-hydroxy-7- methoxybenzofuran) and (5-acety1-7-(4'-bromo)benzyloxy-6-hydroxy-4- methoxybenzofuran), block Kv1.3 with EC50 values of 480 and 400 nM, resp. Both compds. exhibit moderate selectivity over other Kv1-family channels and HERG, are not cytotoxic, and suppress human T cell proliferation at low micromolar concens.

cell proliferation at low micromolar concis

IT 880479-06-1P

ACCESSION NUMBER:

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and voltage-gated potassium channel activity of khellinone analogs)

RN 880479-06-1 CAPLUS

CN Ethanone, 1,1'-[1,6-hexanediylbis[oxy(4-ethoxy-7-methoxy-6,5-

benzofurandiyl)]]bis- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 46 THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

2004:446728 CAPLUS Full-text

L15 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN

DOCUMENT NUMBER: 141:251728

TITLE: 1,6-Bis(5-acetyl-4,7-dimethoxybenzofuran-6-

vloxy)hexane

AUTHOR(S): Baell, Jonathan B.; Gable, Robert W.; Harvey, Andrew

J.

CORPORATE SOURCE: Structural Biology Chemistry Group, The Walter and

Eliza Hall Institute of Medical Research,

Biotechnology Centre, Bundoora, Victoria, 3086,

Australia

SOURCE: Acta Crystallographica, Section E: Structure Reports

Online (2004), E60(6), o996-o997 CODEN: ACSEBH: ISSN: 1600-5368

PUBLISHER: International Union of Crystallography

DOCUMENT TYPE: Journal; (online computer file)

LANGUAGE: English

AB The khellinone dimer, 1,6-bis-(5-acetyl-4,7-dimethoxybenzofuran-6-yloxy)hexane, C30H34010, was prepared as part of Kv1.3 ion channel blockers. Crystallog. data are given. The dimer lies on a center of symmetry, and adopts an extended structure such that the separation between the benzofuran groups is 9.927(3) Å. C-H···O H bonds link the mols. into linear chains which lie parallel to the [201] direction.

IT 605665-31-4P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and crystal structure of)

RN 605665-31-4 CAPLUS

CN Ethanone, 1,1'-[1,6-hexanediylbis[oxy(4,7-dimethoxy-6,5-benzofurandiyl)]]bis- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L15 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2004:178993 CAPLUS Full-text

DOCUMENT NUMBER: 140:375004

TITLE: Khellinone Derivatives as Blockers of the

Voltage-Gated Potassium Channel Kv1.3: Synthesis and

Immunosuppressive Activity

AUTHOR(S): Baell, Jonathan B.; Gable, Robert W.; Harvey, Andrew

J.; Toovey, Nathan; Herzog, Tanja; Haensel, Wolfram;

Wulff, Heike

CORPORATE SOURCE: Walter and Eliza Hall Institute of Medical Research
Biotechnology Centre, Bundoora, 3086, Australia
SOURCE: Journal of Medicinal Chemistry (2004), 47(9),

2326-2336

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal

LANGUAGE: English
OTHER SOURCE(S): CASREACT 140:375004

AB The voltage-gated potassium channel Kv1.3 constitutes a promising new target for the treatment of T-cell-mediated autoimmune diseases such as multiple sclerosis. In this study, we report the discovery of two new classes of Kv1.3 blockers based on the naturally occurring compound khellinone, 5-acetyl-4,7dimethoxy-6-hydroxybenzofuran: (1) khellinone dimers linked via the alkylation of the 6-hydroxy groups and (2) chalcone derivs. of khellinone formed by Claisen-Schmidt condensation of the 5-acetyl group with aryl aldehydes. In particular, the chalcone 3-(4,7-dimethoxy-6-hydroxybenzofuran-5-yl)-1-phenyl-3-oxopropene and several of its derivs, inhibited Kv1.3 with Kd values of 300-800 nM and a Hill coefficient of 2, displayed moderate selectivity over other Kv1-family K+ channels, suppressed T-lymphocyte proliferation at submicromolar concns., and showed no signs of acute toxicity in mice. Because of their relatively low mol. weight and lipophilicity and their high affinity to Kv1.3, aryl-substituted khellinone derivs, represent attractive lead compds, for the development of more potent and selective Kv1.3 blocking immunosuppressants.

- IT 605665-30-3P 605665-31-4P 605665-32-5P 684278-39-5P
 - RL: BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
- (preparation and immunosuppressive activity of khellinone derivs.)
- RN 605665-30-3 CAPLUS
- CN Ethanone, 1,1'-[1,5-pentanediylbis[oxy(4,7-dimethoxy-6,5-benzofurandiyl)]]bis-(9CI) (CA INDEX NAME)

- RN 605665-31-4 CAPLUS
- CN Ethanone, 1,1'-[1,6-hexanediylbis[oxy(4,7-dimethoxy-6,5-benzofurandiyl)]]bis- (9CI) (CA INDEX NAME)

- RN 605665-32-5 CAPLUS
- CN Ethanone, 1,1'-[1,4-butanediylbis[oxy(4,7-dimethoxy-6,5benzofurandiyl)]]bis- (9CI) (CA INDEX NAME)

- RN 684278-39-5 CAPLUS
- CN Ethanone, 1,1'-[1,7-heptanediylbis[oxy(4,7-dimethoxy-6,5-benzofurandiyl)]]bis- (9CI) (CA INDEX NAME)

L15 ANSWER 4 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2003:757693 CAPLUS Full-text DOCUMENT NUMBER: 139:276752

TITLE: Preparation of divalent ligands based on khellinone derivatives as therapeutic ion channel blocking agents INVENTOR(S): Baell, Jonathan B.; Wulff, Heike; Harvey, Andrew J.;

Norton, Raymond S.; Chandy, George K.

PATENT ASSIGNEE(S): The Walter and Eliza Hall Institute of Medical

Research, Australia SOURCE: PCT Int. Appl., 64 pp.

CODEN: PIXXD2

Patent DOCUMENT TYPE: LANGUAGE: English

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WO					A1		20030925		WO 2003-AU351					20030320				
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB	, BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,	
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC	, EE,	ES,	FI,	GB,	GD,	GE,	GH,	
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE	, KG,	KP,	KR,	KZ,	LC,	LK,	LR,	
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN	, MW,	MX,	MZ,	NI,	NO,	NZ,	OM,	
		PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG	, SK,	SL,	TJ,	TM,	TN,	TR,	TT,	
		TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA	, ZM,	ZW						
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ	, TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,	
		KG,	ΚZ,	MD,	RU,	ТJ,	TM,	AT,	BE,	BG	, CH,	CY,	CZ,	DE,	DK,	EE,	ES,	
		FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC	, NL,	PT,	RO,	SE,	SI,	SK,	TR,	
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ	, GW,	ML,	MR,	NE,	SN,	TD,	TG	
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									AU 2003-212101									
EP	1490349			A1		2004			EP 2003-707912				20030320					
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	PT,	
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL	, TR,	BG,	CZ,	EE,	HU,	SK		
	CN 1656087									CN 2003-811505								
	JP 2005525384																	
	IN 2004DN02795									IN 2004-DN2795								
US	US 2005261301				A1		20051124			US	2005-	005-507925			2	20050705		
PRIORIT:	RIORITY APPLN. INFO.:									ΑU	2002-	1272						
										WO	2003-	AU35	1		W 2	0030	320	
OTHER S	THER SOURCE(S):				MARPAT		139:	2767	52									

GT

AB The title compds. [I; R1-R4 = H, OH, alkyl, alkoxy, etc.; X = a divalent spacer group that provides a spacing between the two aromatic rings to which it is joined of from 6 to 11 atoms when measured across the shortest route between the two aromatic rings; A, B = fused rings independently selected from (un) substituted 5-7 membered (hetero) aromatic and non-aromatic heterocyclic rings; R5, R6 = COR7, C(NR7)R7, CSR7 (R7 = H, alkyl, alkoxy, OH); with the proviso] which can be useful in the modulation of potassium channel activity in cells, including among others Kv1.3 channels found in T-cells, were prepared Thus, reacting khellinone with 1,5-dibromopentane in the presence of cesium carbonate in DMF afforded 65% II which showed Kd of 0.82 µM (Kv1.3) and Kd of 1.5 μM (Kv1.2). The compds. I may also be useful in the treatment or prevention of autoimmune and inflammatory diseases, including multiple sclerosis. Pharmaceutical composition comprising the compound I was claimed. IΤ 605665-30-3P 605665-31-4P 605665-32-5P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of divalent ligands based on khellinone derivs. as therapeutic ion channel blocking agents)

RN 605665-30-3 CAPLUS

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RN

Ethanone, 1,1'-[1,5-pentanediylbis[oxy(4,7-dimethoxy-6,5-benzofurandivl)]]bis- (9CI) (CA INDEX NAME)

605665-31-4 CAPLUS

CN Ethanone, 1,1'-[1,6-hexanediylbis[oxy(4,7-dimethoxy-6,5-benzofurandiyl)]]bis- (9CI) (CA INDEX NAME)

RN 605665-32-5 CAPLUS

CN Ethanone, 1,1'-[1,4-butanediylbis[oxy(4,7-dimethoxy-6,5benzofurandiyl)]]bis- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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